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A BUILDING BLOCK FORMING A C-C OR C-HETERO ATOM BOND UPON RE-ACTION

Technical Field of the Invention S

fer the functional entity precursor with an adjustable efficiency to a recipient reactive ment and a precursor for a functional entity. The building block is designed to transment associated with the reactive group. The invention also relates to a method for group upon recognition between the complementing element and an encoding ele-The present invention relates to a building block comprising a complementing eleransferring a functional entity precursor to recipient a reactive group.

Background

tyl group from 3'-O-acetyladenosine to the 5'-OH of adenosine. The reverse transfer, Acta, 1971, 228,536-543) used a poly(U) template to catalyse the transfer of an ace-.e. the transfer of the acetyl group from a 5'-O-acetyladenosine to a 3'-OH group of The transfer of a chemical entity from one mono-, di- or oligonucleotide to another has been considered in the prior art. Thus, N. M. Chung et al. (Biochim. Biophys. another adenosine, was also demonstrated. 5 8

cedure for peptide synthesis. The synthesis involves the transfer of nascent immobiwhich in turn results in an acyl transfer. It is suggested to attach the amino acid pre-Walder et al. Proc. Natl. Acad. Sci. USA, 1979, 76, 51-55 suggest a synthetic proized polypeptide attached to an oligonucleotide strand to a precursor amino acid attached to an oligonucleotide. The transfer comprises the chemical attack of the amino group of the amino acid precursor on the substitution labile peptidyl ester, cursor to the 5' end of an oligonucleotide with a thiol ester linkage.

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activated thioester is reacted with a first oligo, which is 5'-thiol-terminated, resulting disclosed in Bruick RK et al. Chemistry & Biology, 1996, 3:49-56. The carboxy terin the formation of a thio-ester linked intermediate. The first oligonucleotide and a transformed to an activated thioester upon incubation with Ellman's reagent. The The transfer of a peptide from one oligonucleotide to another using a template is minal of the peptide is initially converted to a thioester group and subsequently 8 ઝ

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(57) Abstract: A building block having the dual capabilities of transferring genetic information and functional entity precursor to a complexes, wherein the complex comprises an encoded molecule linked to an encoding element. Libraries of complexes are useful recipient reactive group is disclosed. The building block may be used in the generation of a single complex or libraries of different (54) Title: A BUILDING BLOCK FORMING A C-C OR A C-HETTERO ATOM BOND UPONREACTION 977870/E0 OW

in the quest for pharmaceutically active compounds.

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second oligonucleotide having a 3' amino group is aligned on a template such that the thioester group and the amino group are positioned in close proximity and a transfer is effected resulting in a coupling of the peptide to the second oligonucleotide through an amide bond.

Summary of the Invention

The present invention relates to a building block of the general formula:

Complementing Element – Linker – Carrier – C-F-connecting group - Functional entity precursor

capable of transferring a Functional entity precursor to a recipient reactive group,

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Complementing Element is a group identifying the Functional entity precursor, Linker is a chemical moiety comprising a spacer and a S-C-connecting

group, wherein the spacer is a valence bond or a group distancing the Functional entity precursor to be transferred from the complementing element and the S-C-connecting group connects the spacer with the Carrier

٠ ئ Carrier is arylene, heteroarylene, C_1 - C_6 alkylene, C_1 - C_6 alkenylene, C_1 - C_6 alkynylene, or - $(CF_2)_{nr}$ - substituted with 0-3 R¹ wherein m is an integer between 1 and

R¹ are independently selected from -H, -OR², -NR²₂, -Halogen, -NO₂, -CN, -C(Halogen)₃, -C(O)R², -C(O)NHR², C(O)NR²₂, -NC(O)R², -S(O)₂NHR², -S(O)₂NR²₂, -S(O)₂R², -P(O)²-R², -S(O)-R², P(O)-OR², -S(O)-OR², -N'R²₃, wherein R² is H, C₁-C₀ alkyl, C₂-C₀ alkenyl, C₂-C₀ alkynyl, or aryl,

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C-F-connecting group is chosen from the group consisting of –SO₂-O-, 25 -O-SO₂-O-, -C(O)-O-, -S'(R³RRrr)-, -C-U-C(V)-O-, -P*(W)_Z-O-, -P(W)-O- where U is -C(R³)_Z-, -NR²- or –O-; V is =O or =NR² and W is -OR² or –N(R²)₂

Functional entity precursor is -C(H)(R³)-R⁴ or functional entity precursor is heteroaryl or aryl optionally substituted with one or more substituents belonging to the group comprising R³ and R⁴.

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Wherein R³ and R⁴ independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cydoalkyl, cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents selected from the group consisting of SnR⁵R³R³, Sn(OR³)R²R³. Sn(OR³)R², BR⁵R³, B(OR⁵)R³, B(OR⁵)R°, B(OR°)R°, B(OR°)R°,

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WO 03/078446

PCT/DK03/00176

OR⁵, OC(=O)R⁵, OC(=O)OR⁵, OC(=O)NR⁵R⁶, SR⁵, S(=O)R⁵, S(=O)₂NR⁵, S(=O)₂NR⁵, NR⁵CR⁵, NR⁵CR⁵, NR⁵CR⁶, NR⁵CR⁶, NR⁵CR⁶, NR⁵CR⁶, NR⁵C(=O)R⁶, NR⁵C(=O)OR⁶, NR⁵C(=O)OR⁶, NR⁵C(=O)NR⁵C, C(=O)NR⁵C, C(=O)NR⁵CC, C(=O)NC⁵CC, C(=O

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R⁵, R⁶, and R⁷ independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl, cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents selected from the group consisting of halogen, CN, CNO, C(halogen)₃₀, =O, OR⁶, OC(=O)R⁶, OC(=O)OR⁶, OC(=O)R⁶, SC(=O)R⁶, SC(=O

S(=O)₂NR²R², NO₂, N₃, NR²R³, N²R³R¹⁰, NR⁵OR², NR⁵R²R², NR²C(=O)R², NR³C(=O)OR², NR³C(=O)NR³R³, C(=O)(OR³)OR³, P²R²R³R³, C(=O)R³, C(=NR³)R³, C(=NOR³)R³, C(=NNR³R³, C(=O)NR³R³, C(=O)NR³R³, C(=O)NR³CR³ or C(=O)NR³R³, wherein R⁵ and R⁵ may together form a 3-8 membered heterocyclic ring or R⁵ and R⁷ may together form a 3-8 membered heterocyclic ring or R³ and R⁷ may together form a 3-8 membered heterocyclic ring or R³ and R⁷ may together form a 3-8 membered heterocyclic ring.

wherein,

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R°, R°, and R¹º Independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl, cycloheteroalkyl, aryl or heteroaryl and wherein R° and R° may together form a 3-8 membered heterocyclic ring or R° and R¹º may together form a 3-8 membered heterocyclic ring or R° and R¹º may together form a 3-8 membered heterocyclic ring.

In the present description and claims, the direction of connections between the various components of a building block should be read left to right. For example an S-C-connecting group –C(=O)-NH- is connected to a Spacer through the carbon atom on the left and to a Carrier through the nitrogen atom on the right hand side.

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The term "C₂-C₇ cycloheteroalkyl" as used herein refers to a radical of totally saturated heterocycle like a cyclic hydrocarbon containing one or more heteroatoms selected from nitrogen, oxygen, phosphor, boron and sulphur independently in the cycle such as pyrrollidine (1- pyrrollidine; 2- pyrrollidine; 3- pyrrollidine; 4- pyrrollidine; 5- pyrrollidine; 5- pyrrazollidine; 3- pyrazollidine; 5- pyrazollidine; 2- imidazollidine; 2- imida-

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zolidine; 3- thiazolidine; 4- thiazolidine; 5- thiazolidine); piperidine (1- piperidine; 2piperidine; 3- piperidine; 4- piperidine; 5- piperidine; 6- piperidine); piperazine (1zolidine; 3- imidazolidine; 4- imidazolidine; 5- imidazolidine); thiazolidine (2- thiapiperazine; 2- piperazine; 3- piperazine; 4- piperazine; 5- piperazine; 6-

pholine; 6- morpholine); thiomorpholine (2- thiomorpholine; 3- thiomorpholine; 4piperazine); morpholine (2- morpholine; 3- morpholine; 4- morpholine; 5- morthiomorpholine; 5- thiomorpholine; 6- thiomorpholine); 1,2-oxathiolane (3-(1,2oxathiolane); 4-(1,2-oxathiolane); 5-(1,2-oxathiolane); 1,3-dioxolane (2-(1,3dioxolane); 4-(1,3-dioxolane); 5-(1,3-dioxolane); tetrahydropyrane; (2-

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tetrahydropyrane; 3-tetrahydropyrane; 4-tetrahydropyrane; 5-tetrahydropyrane; 6-(hexahydropyridazine); 3-(hexahydropyridazine); 4-(hexahydropyridazine); 5-(hexahydropyridazine); 6-(hexahydropyridazine)), [1,3,2]dioxaborolane, tetrahydropyrane); hexahydropyridazine (1-(hexahydropyridazine); 2-[1,3,6,2]dioxazaborocane

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bon atoms. Any is also intended to include the partially hydrogenated derivatives of the carbocyclic systems as well as up to four fused fused aromatic- or partially hy-The term "aryl" as used herein includes carbocyclic aromatic ring systems of 5-7 cardrogenated rings, each ring comprising 5-7 carbon atoms.

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from nitrogen, oxygen and sulphur such as furyl, thienyl, рултоlyl, heteroaryl is also The term "heteroary!" as used herein includes heterocyclic unsaturated ring systems intended to include the partially hydrogenated derivatives of the heterocyclic syscontaining, in addition to 2-18 carbon atoms, one or more heteroatoms selected tems enumerated below.

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The terms "ary!" and "heteroary!" as used herein refers to an aryl which can be opanthracenyl, 2-anthracenyl, 3-anthracenyl), thiophenyl (2-thienyl, 3-thienyl), furyl (2-furyl, 3-furyl), indolyl, oxadiazolyl, isoxazolyl, quinazolinyl, fluorenyl, xanthenyl, zolyl (1,2,3-triazol-1-yl, 1,2,3-triazol-2-yl 1,2,3-triazol-4-yl, 1,2,4-triazol-3-yl), oxapyrazolyl), imidazolyl (1-imidazolyl, 2-imidazolyl, 4-imidazolyl, 5-imidazolyl), triapyrimidinyl, 5-pyrimidinyl, 6-pyrimidinyl), pyrazinyl, pyridazinyl (3- pyridazinyl, 4tionally substituted or a heteroaryl which can be optionally substituted and inthiazolyl), pyridyl (2-pyridyl, 3-pyridyl, 4-pyridyl), pyrimidinyl (2-pyrimidinyl, 4zolyl (2-oxazolyl, 4-oxazolyl, 5-oxazolyl), thiazolyl (2-thiazolyl, 4-thiazolyl, 5isoIndanyl, benzhydryl, acridinyl, thiazolyl, pyrrolyl (2-pyrrolyl), pyrazolyl (3hydroxytetrazolyl, N-hydroxytriazolyl, N-hydroxyimidazolyl, anthracenyl (1cludes phenyl, biphenyl, indenyl, naphthyl (1-naphthyl, 2-naphthyl), N-

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WO 03/078446

PCT/DK03/00176

pyridazinyi, 5-pyridazinyi), quinolyi (2-quinolyi, 3-quinolyi, 4-quinolyi, 5-quinolyi, 6quinolyl, 7-qulnolyl, 8-quinolyl), isoquinolyl (1-isoquinolyl, 3-isoquinolyl, 4isoquinolyl, 5-isoquinolyl, 6-isoquinolyl, 7-isoquinolyl, 8-isoquinolyl),

benzo[bʃfuranyl, 6-benzo[bʃfuranyl, 7-benzo[bʃfuranyl), 2,3-dihydro-benzo[bʃfuranyl oenzo[bʃturanyl), 5-(2,3-dihydro-benzo[bʃturanyl), 6-(2,3-dihydro-benzo[bʃturanyl), 2-(2,3-dihydro-benzo[b]furanyl), 3-(2,3-dihydro-benzo[b]furanyl), 4-(2,3-dihydror-(2,3-dihydro-benzo[b]furanyl), benzo[b]thiophenyl (2-benzo[b]thiophenyl, 3benzo[b]furanyl (2-benzo[b]furanyl, 3-benzo[b]furanyl, 4-benzo[b]furanyl, 5benzo[b]thiophenyl, 4-benzo[b]thiophenyl, 5-benzo[b]thiophenyl, 6-

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benzo[b]thiophenyl, 7-benzo[b]thiophenyl), 2,3-dihydro-benzo[b]thiophenyl (2-(2,3indolyl, 3-indolyl, 4-indolyl, 5-indolyl, 6-indolyl, 7-indolyl), indazole (1-indazolyl, 3-Jihydro-benzo[b]thiophenyl), 3-(2,3-dihydro-benzo[b]thiophenyl), 4-(2,3-dihydroindazolyl, 4-indazolyl, 5-indazolyl, 6-indazolyl, 7-indazolyl), benzimidazolyl (1benzo[b]thiophenyl), 7-(2,3-dihydro-benzo[b]thiophenyl), indolyl (1-indolyl, 2benzo[b]thiophenyl), 5-(2,3-dihydro-benzo[b]thiophenyl), 6-(2,3-dihydro-

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senzoxazolyl, 2-benzoxazolyl), benzothiazolyl (1-benzothiazolyl, 2-benzothiazolyl, 4-benzothiazolyl, 5-benzothiazolyl, 6-benzothiazolyl, 7-benzothiazolyl), carbazolyl benzimidazolyl, 2-benzimidazolyl, 4-benzimidazolyl, 5-benzimidazolyl, 6benzimidazolyl, 7-benzimidazolyl, 8-benzimidazolyl), benzoxazolyl (1-

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(1-carbazolyl, 2-carbazolyl, 3-carbazolyl, 4-carbazolyl), 5H-dibenz[b,f]azepine (5Hdibenz[b,f]azepin-1-yl, 5H-dibenz[b,f]azepine-2-yl, 5H-dibenz[b,f]azepine-3-yl, 5Hdibenz[b,f]azepine-2-yl, 10,11-dihydro-5H-dibenz[b,f]azepine-3-yl, 10,11-dihydrodibenz[b,f]azepine (10,11-dihydro-5H-dibenz[b,f]azepine-1-yl, 10,11-dihydro-5Hdibenz[b,f]azepine-4-yl, 5H-dibenz[b,f]azepine-5-yl), 10,11-dihydro-5H-5H-dibenz[b,f]azepine-4-yl, 10,11-dihydro-5H-dibenz[b,f]azepine-5-yl).

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library. Interaction with host molecules like enzymes, receptors and polymers is typically mediated through van der waal's interactions, polar- and ionic interactions and building blocks and during library synthesis. Analogously, reactive elements may be The Functional Entity carries elements used to interact with host molecules and oppi-stacking effects. Substituents mediating said effects may be masked by methods ionally reactive elements allowing further elaboration of an encoded molecule of a known to an individual skilled in the art (Greene, T. W.; Wuts, P. G. M. Protective avoid undesired interactions or reactions during the preparation of the individual Groups in Organic Synthesis; 3rd ed.; John Wiley & Sons: New York, 1999.) to

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PCT/DK03/00176

masked by suitably selected protection groups. It is appreciated by one skilled in the art that by suitable protection, a functional entity may carry a wide range of substi-

Entity may be revealed by un-masking allowing further synthetic operations. Finally, The Functional Entity Precursor is a masked Functional Entity that is incorporated nto an encoded molecule. After incorporation, reactive elements of the Functional elements mediating recognition of host molecules may be un-masked.

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In a certain aspect of the invention, Functional entity precursor is -C(H)(R^{11})- R^{11} . or functional entity precursor is heteroaryl or aryl substituted with 0-3 $\mathrm{R}^{1},$ 0-3 R^{13} and 0-3 R¹⁶, wherein 우

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 R^{11} and R^{11} are independently H, or selected among the group consisting of a $C_1\text{-}C_8$ heteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R^{12} , 0-3 R^{13} alkyl, Cz-Ce alkenyl, Cz-Ce alkynyl, C₄-Ce alkadienyl, C₃-Cγ cydoalkyl, C₃-Cγ cydlo-

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or R11 and R115 are C1-C3 alkylene-NR12, C1-C3 alkylene-NR12C(O)R16, C1-C3 altered by the constant of the con kylene-NR 12 C(O)OR 19 , C,-C $_2$ alkylene-O-NR 12 , C,-C $_2$ alkylene-O-NR 12 C(O)R 16 , and 0-3 R¹⁵.

where R^{12} is H or selected independently among the group consisting of $\mathsf{C}_1\text{-}\mathsf{C}_8$ alkyi, C_z-C₈ alkenyi, C_z-C₈ alkynyi, C₃-C, cycloalkyi, C₃-C, cycloheteroalkyi, aryi, C₁-C₂ alkylene-O-NR¹²C(O)OR¹⁶ substituted with 0-3 R¹⁵,

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 C_z – C_e alkenyl, C_z – C_e alkynyl, G_z – C_e alkadienyl said group being substituted with 0-2 -NHNHR'7, -C(O)R'7, -SnR¹73, -B(OR'7)2, -P(O)(OR'7)2 or the group consisting of R¹³ is selected independently from -N₃, -CNO, -C(NOH)NH₂, -NHOH, heteroaryl, said group being substituted with 0-3 \mbox{R}^{19} and 0-3 $\mbox{R}^{15},$

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where R^{14} is independently selected from $-NO_{2},\,-C(O)OR^{17},\,-COR^{17},\,-CN,$ -OSiR¹⁷s, -OR¹⁷ and -NR¹⁷z;

alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -- $-NR^{17}-C(O)OR^{10}, -SR^{17}, -S(O)R^{17}, -S(O)_2R^{17}, -COOR^{17}, -C(O)NR^{17}, \text{ and } -S(O)_2NR^{17}, -C(O)NR^{17}, -$ R¹⁹ is H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, aryl or C₁-C₆ R¹⁵ is =0, -F, -CI, -Br, -I, -CN, -NO₂, -OR¹⁷, -NR¹⁷, -NR¹⁷-C(O)R¹⁶, NO2, -R2, -OR2, -SIR3;

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WO 03/078446

PCT/DK03/00176

R¹⁷ is selected independently from H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, aryl, C₁-C₈ al-

 $(\text{Niene-ary}), \text{ or } \text{ Un } \text{ or } \text{ G is H or } \text{C}_1\text{-C}_6 \text{ alkyl and n is } 1,2,3 \text{ or } 4.$

the nature of the complementing element, to the conditions under which the transfer justed in response to the chemical composition of the functional entity precursor, to The function of the carrier is to ensure the transferability of the functional entity precursor. To adjust the transferability a skilled chemist can design suitable substitutions of the carrier by evaluation of initial attempts. The transferability may be adand recognition is performed, etc. S

such compounds a broad range of recipient reactive groups may be employed in the tween 1 and 10, and C-F-connecting group is $-SO_z$ -O-. Due to the high reactivity of in a preferred embodiment, the carrier is selected from the group consisting of arylene, heteroarylene or -(CF₂) $_{\!\!\!m}$ substituted with 0-3 R¹ wherein m is an integer belene, construction of carbon-carbon bonds or carbon-hetero atom bonds. . 5

in another preferred embodiment of the invention, the carrier is -(CF_2) $_{\text{m}^-}$ wherein m is an integer between 1 and 10, the C-F-connecting group is –SO $_{2}$ -O-; and the functional entity precursor is aryl or heteroaryl substituted with 0-3 \mbox{R}^{11} , 0-3 \mbox{R}^{13} and 0-3

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The C-F-connecting group determines in concert with the carrier the transferability of the functional entity precursor. In a preferred embodiment, the C-F-connecting

group is -S*(R¹¹)-,

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group consisting of -SO₂-O-, and -S*(R¹⁷)-; wherein R¹⁷ is selected independently In another preferred embodiment, the C-F-connecting group is chosen from the from H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, aryl, C₁-C₈ alkylene-aryl. In the presence of a catalyst comprising transition metals such as Pd, Ni or Cu, an aromatic moiety may be transferred from the C-F-connecting group to a recipient reactive group. Further, the transfer may be initiated by adding the catalyst, independently of the annealing of encoding - and complementing elements.

PCT/DK03/00176

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The S-C-connecting group provide a means for connecting the Spacer and the Carrier. As such it is primarily of synthetic convenience and does not influence the function of a building block.

The spacer serves to distance the functional entity precursor to be transferred from

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the bulky complementing element. Thus, when present, the identity of the spacer is not crucial for the function of the building block. It may be desired to have a spacer which can be cleaved by light. In this case, the spacer is provided with e.g. the

group

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In the event an increased hydrophilicity is desired the spacer may be provided with a polyethylene glycol part of the general formula:

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In a preferred embodiment, the complementing element serves the function of transferring genetic information e.g. by recognising a coding element. The recognition implies that the two parts are capable of interacting in order to assemble a complementing element – coding element complex. In the biotechnological field a variety of interacting molecular parts are known which can be used according to the invention. Examples include, but are not restricted to protein-protein interactions, protein-polysaccharide interactions, RNA-protein interactions, DNA-DNA interactions, DNA-RNA interactions, biotin-streptavidin interactions, enzymeligand interactions, antibody-ligand interaction, protein-ligand interaction, etc.

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The interaction between the complementing element and coding element may result in a strong or a weak bonding. If a covalent bond is formed between the parties of the affinity pair the binding between the parts can be regarded as strong, whereas the establishment of hydrogen bondings, interactions between hydrophobic do-

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WO 03/078446

PCT/DK03/00176

mains, and metal chelation in general results in weaker bonding. In general relatively weak bonding is preferred. In a preferred aspect of the invention, the complementing element is capable of reversible interacting with the coding element so as to provide for an attachment or detachment of the parts in accordance with the chang-

5 ing conditions of the media.

In a preferred aspect of the invention, the interaction is based on nucleotides, i.e. the complementing element is a nucleic acid. Preferably, the complementing element is a sequence of nucleotides and the coding element is a sequence of nucleotides and the coding element. The sequence of nucleotides carries a series of nucleobases on a backbone. The nucleobases may be any chemical entity able to be specifically recognized by a complementing entity. The nucleobases are usually selected from the natural nucleobases (adenline, guanline, uracil, thymine, and cytosine) but also the other nucleobases obeying the Watson-Crick hydrogen-bonding rules may be used, such as the synthetic nucleobases disclosed in US 6,037,120. Examples of natural and non-natural nucleobases able to perform a specific pairing are shown in figure 2. The backbone of the sequence of nucleotides may be any backbone able to aggregate the nucleobases is a sequence. Examples of backbones are shown in figure 4. In some aspects of the invention the addition of non-specific nucleobases to the complementing element is

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The coding element can be an oligonucleotide having nucleobases which complements and is specifically recognised by the complementing element, i.e. in the event the complementing element contains cytosine, the coding element part contains guanine and visa versa, and in the event the complementing element contains thymine or uracil the coding element contains adenine.

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The complementing element may be a single nucleobase. In the generation of a library, this will allow for the incorporation of four different functional entities into the template-directed molecule. However, to obtain a higher diversity a complementing element preferred at least three nucleositides. Theoretically, this will provide for 4² and 4³, respectively, different functional entities uniquely identified by the complementing element. The complementing element will usually not comprise more than 100 nucleotides. It is preferred to have complementing elements with a sequence of 3 to 30 nucleotides.

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The building blocks of the present invention can be used in a method for transferring a functional entity precursor to a recipient reactive group, said method comprising the steps of

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providing one or more building blocks as described above and contacting the one or more building blocks with a corresponding encoding element associated with a recipient reactive group under conditions which allow for a recognition between the one or more complementing elements and the encoding elements, said contacting being performed prior to, simultaneously with, or subsequent to a transfer of the functional entity precursor to the recipient reactive group.

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The encoding element may comprise one, two, three or more codons, i.e. sequences that may be specifically recognised by a complementing element. Each of quences that may be separated by a suitable spacer group. Preferably, all or at least a majority of the codons of the template are arranged in sequence and each of the codons are separated from a neighbouring codon by a spacer group. Generally, it is preferred to have more than two codons on the template to allow for the synthesis of more complex encoded molecules. In a preferred aspect of the invention the number of codons of the encoding element is 2 to 100. Still more preferred are encoding elements comprising 3 to 10 codons. In another aspect, a codon comprises 1 to 50 nucleotides and the complementing element comprises a sequence of nucleotides complementary to one or more of the encoding sequences.

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The recipient reactive group may be associated with the encoding element in any appropriate way. Thus, the reactive group may be associated covalently or non-covalently to the encoding element. In one embodiment the recipient reactive group is linked covalently to the encoding element through a suitable linker which may be separately cleavable to release the reaction product. In another embodiment, the reactive group is coupled to a complementing element, which is capable of recognising a sequence of nucleotides on the encoding element, whereby the recipient reactive group becomes attached to the encoding element by hybridisation. Also, the recipient reactive group may be part of a chemical scaffold, i.e. a chemical entity having one or more reactive groups available for receiving a functional entity precursor from a building block.

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WO 03/078446

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The recipient reactive group may be any group able to participate in cleaving the bond between the carrier and the functional entity precursor to release the functional entity precursor. Typically, the recipient reactive group is a nucleophilic atom such as S, N, O, C or P. Scheme 1a shows the transfer of an alkyl group and scheme 1b

5 shows the transfer of an vinyl group...

Scheme 1a



Scheme 1b

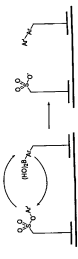
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CN, COOR, COR, NO2, SO2R, S(=O)R, SO2NR2, F Course. Nitrogen - Sultir- and Carbon Nucleoph Alternatively, the recipient reactive group is a organometallic compound as shown in scheme 2.

Scheme 2

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According to a preferred aspect of the invention the building blocks are used for the formation of a library of compounds. The complementing element of the building block is used to identify the functional entity. Due to the enhanced proximity between reactive groups when the complementing entity and the encoding element are

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menting element having a unique sequence of nucleotides, which identifies the funcfunctional entity. The unique identification of the functional entity enable the possibildetermined. Also the sequence of reaction and the type of reaction involved can be ferred to a scaffold, not only the identity of the transferred functional entities can be the complementing element is transferred to the encoding element associated with recipient reactive group. Thus, it is preferred that the sequence of the complementing element is unique in the sense that the same sequence is not used for another ity of decoding the encoding element in order to determine the synthetic history of the molecule formed. In the event two or more functional entities have been transdetermined by decoding the encoding element. Thus, according to a preferred embodiment of the invention, each different member of a library comprises a complecontacted, the functional entity precursor together with the identity programmed in

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Brief description of the drawings 5

Figure 1. Two setups for Functional Entity Transfer Figure 2. Examples of specific base pairing

Figure 3. Example of non-specific base-pairing

Figure 4. Backbone examples

Figure 5 Three examples of building blocks

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Detailed Description of the Invention

A building block of the present invention is characterized by its ability to transfer its new covalent bond between the recipient reactive group and cleaving the bond befunctional entity precursor to a recipient reactive group. This is done by forming a ween the carrier moiety and the functional entity precursor of the building block.

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between functional entity precursor 1 and 2 forming a covalent bond between these concurrent with the cleavage of the bond between functional entity precursor 2 and Iwo setups for generalized functional entity precursor transfer from a building block are depicted in figure 1. In the first example, one complementing element of a buildits linker. In the second example, a template brings together two building blocks reng block recognizes a coding element carrying another functional entity precursor, nence bringing the functional entities in close proximity. This results in a reaction sulting in functional entity precursor transfer from one building block to the other.

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WO 03/078446

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PCT/DK03/00176

compound is an example of a building block wherein the linker is backbone attached at the 3'-position. The first part of the linker, i.e. the spacer, is an aliphatic chain end-Figure 5 illustrates three specific compounds according to the invention. For illustracarbonyl group of the S-C-connecting group is a benzene ring holding the C-F Conwhich is an N-acylated arylmethyleamine. The carrier attached to the left hand side tive purposes the individual features used in the claims are indicated. The upper ing in a nitrogen atom. The nitrogen atom bridges to the S-C-connecting group, S

necting group in the para position. The C-F Connecting group is a positively charged group, such an amine or a thiol, Functional Entity Precursor is transferred to benzysulfur atom which is attached to the Functional Entity Precursor, in this case a benzy group. When the building block is presented to a nucleophilic recipient reactive ate the recipient reactive group. 2

nucleophile the Functional Entity Precursor is transferred resulting in an alkylation of Through another phosphate group and a PEG linker the complementing element is inked via an amide bond to the Carrier. When the building block is presented to a The middle compound illustrates a 5' attachment of a linker. The linker is linked through a phosphate group and extends into a three membered aliphatic chain. the nucleophile.

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attaches to the 5 position of a pyrimidine type nucleobase and extents through an lphaamino methyl benzoic acid derivative. The functional entity precursor can be transferred to a nucleophilic recipient reactive group e.g. an amine or a thiol forming an he lower compound illustrates a nucleobase attachment of the linker. The linker - ß unsaturated N-methylated amide to the S-C-connecting group, which is a 4allylic amine or thiol.

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-C(H)(R³)-R⁴ or functional entity precursor is heteroaryl or aryl optionally substituted with one or more substituents belonging to the group comprising $\ensuremath{R^3}$ and $\ensuremath{R^4}$. In a According to the invention, the functional entity precursor is of the formula further preferred embodiment, ဓ

kadienyl, Cs-C, cycloalkyl, Cs-C, cycloheteroalkyl, aryl or heteroaryl, optionally sub-R³ and R⁴ independently is H, C₁-C₅ alkyl, C₂-C₅ alkenyl, C₂-C₅ alkynyl, C₄-C₅ alstituted with one or more substituents selected from the group consisting of

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gen, CN, CNO, C(halogen)₃, =0, OR⁵, OC(=0)R⁵, OC(=0)OR⁵, OC(=0)NR⁵R˚, SR⁵, S(=0)R[¢], S(=0)₂R[¢], S(=0)₂NR[¢]R[¢], NO₂, N₃, NR⁶R[¢]R[¢], N'R⁶R⁶R⁷, NR⁵OR⁸, NR⁵NR⁹R⁷, SnR⁵R˚,R′, Sn(OR˚)R°R′, Sn(OR˚)(OR˚)R′, BR⁵R°, B(OR˚)R°, B(OR˚)(OR˚), halo- $\mathsf{NR}^{\mathsf{c}}\mathsf{C}(=\mathsf{O})\mathsf{R}^{\mathsf{g}},\,\mathsf{NR}^{\mathsf{c}}\mathsf{C}(=\mathsf{O})\mathsf{OR}^{\mathsf{g}},\,\mathsf{NR}^{\mathsf{f}}\mathsf{C}(=\mathsf{O})\mathsf{NR}^{\mathsf{g}}\mathsf{R}^{\mathsf{f}},\,\mathsf{NC},\,\mathsf{P}(=\mathsf{O})(\mathsf{OR}^{\mathsf{d}})\mathsf{OR}^{\mathsf{g}},\,\mathsf{P}^{\mathsf{f}}\mathsf{R}^{\mathsf{f}}\mathsf{R}^{\mathsf{f}},$ C(=0)R³, C(=NR⁵)R˚, C(=NOR˚)R˚, C(=NNR⁵R˚), C(=0)OR˚, C(=0)NR⁵R˚, C(=O)NR5OR, C(=O)NR5NR6R', C(=NR5)NR6R', C(=NOR5)NR6R' or R",

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R⁵, R⁶, R⁷ and R⁸ independently is H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₆ together form a 3-8 membered heterocyclic ring or R^{δ} and R^{γ} may together form a 3alkadienyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl or heteroaryl and wherein R^ϵ and R^ϵ may together form a 3-8 membered heterocyclic ring or R^ϵ and R^γ may 8 membered heterocyclic ring,

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in another prefered embodiment,

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aryl or heteroaryl, optionally substituted with one or more substituents selected from R3 and R4 independently is H, C1-C8 alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, OC(=0)0R°, OC(=0)NR°R°, SR°, S(=0)R°, S(=0)2R°, S(=0)2NR°R°, NO2, NR°R°, P(=0)(OR⁵)OR⁶, C(=0)R⁵, C(=NR⁵)R⁸, C(=NOR⁵)R⁸, C(=NNR⁵R⁸), C(=0)OR⁵, the group consisting of halogen, CN, C(halogen)₃, =0, OR^5 , $OC(=O)R^5$, NR°OR°, NR°NR°R', NR°C(=0)R°, NR°C(=0)OR°, NR°C(=0)NR°R',

C(=O)NR\$R\$, C(=O)NR\$OR\$, C(=O)NR\$NR\$R7, C(=NR\$)NR\$R7, C(=NOR\$)NR\$R7 or

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wherein

bered heterocyclic ring or ${
m R}^5$ and ${
m R}^7$ may together form a 3-8 membered heterocyc-R⁵, R⁹, R⁷ and R⁸ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein R5 and R9 may together form a 3-8 memic ring or R⁶ and R⁷ may together form a 3-8 membered heterocyclic ring,

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in still another prefered embodiment,

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aryl or heteroaryl, optionally substituted with one or more substituents selected from R3 and R4 independently is H, C1-C6 alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, OC(=0)NR5R⁸, SR⁵, S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR5R⁶, NO₂, NR5R⁶, NR⁶OR⁶, NR^sNR^sR', NR^sC(=0)R^s, NR^sC(=0)OR^s, NR^sC(=0)NR^sR', P(=0)(OR^s)OR^s, the group consisting of F, Cl, CN, CF_3 , =O, OR^5 , $OC(=O)R^5$, $OC(=O)OR^5$,

C(=0)R°, C(=NR°)R°, C(=NOR°)R°, C(=NNR°FR°), C(=0)OR°, C(=0)NR°FR°,

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 $C(=O)NR^5OR^6$, $C(=O)NR^5NR^6R^7$, $C(=NR^5)NR^9R^7$ or R^8 , $C(=NOR^5)NR^9R^7$ or R^8 ,

bered heterocyclic ring or ${
m R}^5$ and ${
m R}^7$ may together form a 3-8 membered heterocyc-R⁵, R⁹, R⁷ and R⁸ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein R^{δ} and R^{θ} may together form a 3-8 memic ring or R⁸ and R⁷ may together form a 3-8 membered heterocyclic ring, Ŋ

in still another prefered embodiment,

aryl or heteroaryl, optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, =O, OR 6 , S(=O)R 5 , S(=O)₂NR 5 R 6 , R3 and R4 independently is H, C1-C6 alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, NO2, NR⁵R°, NR⁵C(=0)R³, NR⁵C(=0)OR³, NR⁵C(=0)NR³R⁷, C(=0)R⁵, C(=NOR⁸)R⁸, C(=O)OR⁶, C(=O)NR⁵R⁸, C(=O)NR⁵OR⁸ or R⁸, 2

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bered heterocyclic ring or R⁵ and R⁷ may together form a 3-8 membered heterocyc-R⁶, R⁹, R⁷ and R⁸ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein R^5 and R^6 may together form a 3-8 memic ring or R⁶ and R⁷ may together form a 3-8 membered heterocyclic ring,

in still another prefered embodiment,

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phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CFs, NR°C(=0)OR°, NR°C(=0)NR°R′, C(=0)R°, C(=NOR°)R°, C(=0)OR°, C(=0)NR°R°, R3 and R4 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl, morpholinyl, =0, OR⁵, S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR⁵R⁹, NO₂, NR⁵R⁹, NR⁵C(=0)R⁹, C(=O)NR5OR6 or R8,

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wherein,

bered heterocyclic ring or R^{5} and R^{7} may together form a 3-8 membered heterocyc-R⁵, R³, R⁷ and R⁸ independently is H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein R5 and R8 may together form a 3-8 memic ring or R⁸ and R⁷ may together form a 3-8 membered heterocyclic ring, ജ

in still another prefered embodiment, 33

R³ and R⁴ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopenty) or cyclohexyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CFs, =O, OR $^{\circ}$, S(=O)R $^{\circ}$, S(=O) $_{\rm z}$ R $^{\circ}$, S(=0)2NR*R*, NO2, NR*R*, NR*C(=0)R*, NR*C(=0)OR*, NR*C(=0)NR*R7,

.C(=0)R⁵, C(=NOR⁵)R⁶, C(=0)OR⁵, C(=0)NR⁵R⁶, C(=0)NR⁵OR⁶ or R⁸, ß

wherein,

bered heterocyclic ring or R^{5} and R^{7} may together form a 3-8 membered heterocyc-R⁵, R⁶, R⁷ and R⁸ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyi, aryl or heteroaryl and wherein R^{δ} and R^{θ} may together form a 3-8 memlic ring or R° and R7 may together form a 3-8 membered heterocyclic ring,

in still another prefered embodiment,

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pholinyl optionally substituted with one or more substituents selected from the group R³ and R⁴ independently is H, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or mor-NR⁵R°, NR⁵C(=0)R°, NR⁵C(=0)OR°, NR⁵C(=0)NR⁶R⁷, C(=0)R⁵, C(=NOR⁵)R⁶, consisting of F, CI, CN, CF₃, =O, OR⁵, S(=O)R⁵, S(=O)₂NR⁵R⁵, NO₂, C(=0)OR⁵, C(=0)NR⁶R⁸, C(=0)NR⁵OR⁸ or R⁸,

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bered heterocyclic ring or R5 and R7 may together form a 3-8 membered heterocyc-R[¢], R[¢], R⁷ and R⁸ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein R^{δ} and R^{δ} may together form a 3-8 memlic ring or R⁸ and R⁷ may together form a 3-8 membered heterocyclic ring,

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in still another prefered embodiment,

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isoquinolinyl optionally substituted with one or more substituents selected from the R³ and R⁴ independently is H, phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or group consisting of F, Cl, CN, CF₃, =0, OR $^{\circ}$, S(=0)R $^{\circ}$, S(=0)₂NF $^{\circ}$ S(=0)₂NR $^{\circ}$ R $^{\circ}$, NO2, NR*R*, NR*C(=0)R*, NR*C(=0)OR*, NR*C(=0)NR*R7, C(=0)R*, $C(=NOR^5)R^6$, $C(=O)OR^6$, $C(=O)NR^5R^8$, $C(=O)NR^5OR^9$ or R^8 ,

wherein. ജ

bered heterocyclic ring or ${
m R}^5$ and ${
m R}^7$ may together form a 3-8 membered heterocyo-R⁵, R⁵, R⁷ and R⁸ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein R^{δ} and R^{θ} may together form a 3-8 memlic ring or R⁸ and R⁷ may together form a 3-8 membered heterocyclic ring,

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WO 03/078446

PCT/DK03/00176

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in still another prefered embodiment,

NR⁵C(=O)NR⁹R', C(=O)R⁵, C(=NOR⁵)R⁸, C(=O)OR⁵, C(=O)NR⁵R⁸, C(=O)NR⁵OR⁸ R³ and R⁴ independently is H, phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF $_{\rm s}$ =O, OR $^{\rm s}$ S(=0)R°, S(=0)₂R°, S(=0)₂NR°R°, NO₂, NR°R°, NR°C(=0)R°, NR°C(=0)OR°,

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wherein

bered heterocyclic ring or R^{δ} and R^{γ} may together form a 3-8 membered heterocyc-R⁵, R⁸, R⁷ and R⁸ independently is H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and wherein ${\sf R}^5$ and ${\sf R}^6$ may together form a 3-8 memic ring or R⁶ and R⁷ may together form a 3-8 membered heterocyclic ring,

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in still another prefered embodiment,

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tionally substituted with one or more substituents selected from the group consisting NR°C(=0)R°, NR°C(=0)OR°, NR°C(=0)NR°R', C(=0)R°, C(=NOR°)R°, C(=0)OR°, R3 and R4 independently is H, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl opof F, CI, CN, CF₃, =0, OR $^{\circ}$, S(=0)R $^{\circ}$, S(=0) $_{2}$ R $^{\circ}$, S(=0) $_{2}$ NR $^{\circ}$ R $^{\circ}$, NO $_{2}$, NR $^{\circ}$ R $^{\circ}$, C(=0)NR5R6, C(=0)NR5OR6 or R8,

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bered heterocyclic ring or R^{ϵ} and R^{γ} may together form a 3-8 membered heterocyc-Re, Re, R7 and R8 independently is H, C1-Ce alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl or heteroaryl and wherein R^6 and R^θ may together form a 3-8 memic ring or R^{θ} and R^{7} may together form a 3-8 membered heterocyclic ring,

in still another prefered embodiment,

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R³ and R⁴ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl optionally substituted with one or more substituents seected from the group consisting of F, Cl, CN, CFs, =0, OR $^{\circ}$, S(=0)R $^{\circ}$, S(=0) $^{\circ}$ R $^{\circ}$, S(=0)2NR5R°, NO2, NR5R°, NR5C(=0)R°, NR5C(=0)OR°, NR5C(=0)NR8R7, C(=0)R⁵, C(=NOR⁵)R⁸, C(=0)OR⁶, C(=0)NR⁵R⁸, C(=0)NR⁵OR⁸ or R⁸,

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inyl or isoquinolinyl and wherein R^6 and R^6 may together form a 3-8 membered hetcyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quino-R⁵, R³, R⁷ and R⁸ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl,

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erocyolic ring or R5 and R7 may together form a 3-8 membered heterocyolic ring or R⁸ and R⁷ may together form a 3-8 membered heterocyclic ring,

in still another prefered embodiment,

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pholinyl optionally substituted with one or more substituents selected from the group R3 and R4 independently is H, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or mor-NR⁵R°, NR⁵C(=O)R°, NR⁵C(=O)OR°, NR⁵C(=Ò)NRªR7, C(=O)R⁵, C(=NOR⁵)R°, consisting of F, Cl, CN, CF₃, =O, OR⁶, S(=O)R⁶, S(=O)₂R⁶, S(=O)₂NR⁶R⁶, NO₂, $C(=O)OR^5$, $C(=O)NR^5R^6$, $C(=O)NR^5OR^8$ or R^8 ,

wherein. 9

inyl or isoquinolinyl and wherein R⁵ and R⁶ may together form a 3-8 membered heterocyclic ring or R5 and R7 may together form a 3-8 membered heterocyclic ring or cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quino-R5, R6, R7 and R8 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, Re and R7 may together form a 3-8 membered heterocyclic ring,

in still another prefered embodiment,

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isoquinolinyl optionally substituted with one or more substituents selected from the R3 and R4 independently is H, phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or group consisting of F, CI, CN, CF_{31} =0, OR^6 , $S(=O)R^5$, $S(=O)_2R^5$, $S(=O)_2NR^5R^5$, NO2, NR⁵R°, NR⁵C(=0)R°, NR⁵C(=0)OR°, NR⁵C(=0)NR⁶R′, C(=0)R⁶, $C(=NOR^5)R^4$, $C(=O)OR^5$, $C(=O)NR^5R^6$, $C(=O)NR^5OR^9$ or R^3 ,

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linyl or isoquinolinyl and wherein R⁵ and R⁶ may together form a 3-8 membered heterocyclic ring or R^5 and R^7 may together form a 3-8 membered heterocyclic ring or cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quino-R⁵, R⁶, R⁷ and R⁸ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, Re and R7 may together form a 3-8 membered heterocyclic ring,

in still another prefered embodiment, ജ

 $NR^5C(=0)NR^5R^7,\;C(=0)R^5,\;C(=NOR^5)R^8,\;C(=0)OR^5,\;C(=0)NR^5R^8,\;C(=0)NR^5OR^8$ R3 and R4 independently is H, phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF₃, =O, OR⁵, S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR⁵R⁶, NO₂, NR⁵R⁶, NR⁵C(=0)R⁶, NR⁵C(=0)OR⁶,

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WO 03/078446

6

PCT/DK03/00176

wherein.

linyl or isoquinolinyl and wherein R^{ϵ} and R^{θ} may together form a 3-8 membered heterocyclic ring or R5 and R7 may together form a 3-8 membered heterocyclic ring or cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quino-R5, R6, R7 and R8 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl,

in still another prefered embodiment,

Re and R7 may together form a 3-8 membered heterocyclic ring,

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tionally substituted with one or more substituents selected from the group consisting NR⁵C(=0)R⁸, NR⁵C(=0)OR⁸, NR⁵C(=0)NR⁸R⁷, C(=0)R⁵, C(=NOR⁵)R⁸, C(=0)OR⁵, R3 and R4 independently is H, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl opof F, CI, CN, CF₃, =O, OR⁶, S(=O)R⁵, S(=O)₂R⁵, S(=O)₂NR⁵R⁹, NO₂, NR⁵R⁶, C(=O)NR5R°, C(=O)NR5OR° or R°, 9

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linyl or isoquinolinyl and wherein R^{5} and R^{6} may together form a 3-8 membered heterocyclic ring or R⁵ and R⁷ may together form a 3-8 membered heterocyclic ring or cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quino- R^5 , R^7 and R^8 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, Re and R7 may together form a 3-8 membered heterocyclic ring,

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R³ and R⁴ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF3, =0, OR $^{\rm s}$, S(=0)R $^{\rm s}$, S(=O) $_{\rm z}$ R $^{\rm s}$, S(=0)2NR3R8, NO2, NR5R6, NR5C(=0)R8, NR5C(=0)OR8, NR5C(=0)NR8R7, in still another prefered embodiment,

C(=0)R⁵, C(=NOR⁵)R⁶, C(=0)OR⁵, C(=0)NR⁶R⁶, C(=0)NR⁵OR⁶ or R⁸,

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gether form a 3-8 membered heterocyclic ring or ${\rm R}^{\rm s}$ and ${\rm R}^{\rm r}$ may together form a 3-8 R⁵, R⁶, R⁷ and R⁸ independently is H, methyl, ethyl, propyl or butyl and wherein R⁵ and R^{θ} may together form a 3-8 membered heterocyclic ring or R^{δ} and R^{γ} may to-

in still another prefered embodiment,

membered heterocyclic ring,

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pholinyl optionally substituted with one or more substituents selected from the group R³ and R⁴ independently is H, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or mor-છ

NR*R*, NR*C(=0)R*, NR*C(=0)OR*, NR*C(=0)NR*R7, C(=0)R\$, C(=NOR*),R*, consisting of F, CI, CN, CF₃, =O, OR⁶, S(=O)R⁵, S(=O)₂R⁵, S(=O)₂NR⁵R⁶, NO₂, C(=O)OR⁵, C(=O)NR⁶R⁶, C(=O)NR⁵OR⁸ or R⁸,

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- gether form a 3-8 membered heterocyclic ring or $\mbox{R}^{\mbox{\scriptsize a}}$ and $\mbox{R}^{\mbox{\scriptsize 7}}$ may together form a 3-8 R⁶, R⁹, R⁷ and R⁸ independently is H, methyl, ethyl, propyl or butyl and wherein R⁵ and R^{α} may together form a 3-8 membered heterocyclic ring or R^{δ} and R^{γ} may tomembered heterocyclic ring,
- in still another prefered embodiment, 9

isoquinolinyl optionally substituted with one or more substituents selected from the R3 and R4 independently is H, phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or group consisting of F, Cl, CN, CFs, =0, OR⁵, S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR⁵R⁵, NO_2 , NR^5R^6 , $NR^5C(=0)R^8$, $NR^5C(=0)OR^6$, $NR^5C(=0)NR^6R^7$, $C(=0)R^5$,

C(=NOR⁵)R⁹, C(=0)OR⁵, C(=0)NR⁵R⁸, C(=0)NR⁵OR⁸ or R⁸,

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gether form a 3-8 membered heterocyclic ring or ${\sf R}^{\sf a}$ and ${\sf R}^{\sf r}$ may together form a 3-8 R⁶, R⁶, R⁷ and R⁸ independently is H, methyl, ethyl, propyl or butyl and wherein R⁵ and R^a may together form a 3-8 membered heterocyclic ring or R⁵ and R⁷ may tomembered heterocyclic ring,

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in still another prefered embodiment,

NR⁶C(=0)NR⁸R', C(=0)R⁶, C(=NOR⁶)R⁸, C(=0)OR⁶, C(=0)NR⁵OR⁶ R3 and R4 independently is H, phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF3, =O, OR⁵, S(=0)R°, S(=0)2R°, S(=0)2NR5R°, NO2, NR6R°, NR5C(=0)R°, NR5C(=0)OR°,

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wherein, or R°.

gether form a 3-8 membered heterocyclic ring or R^{g} and R^{f} may together form a 3-8 R^s, R°, R7 and R⁸ independently is H, methyl, ethyl, propyl or butyl and wherein R⁵ and R^{θ} may together form a 3-8 membered heterocyclic ring or R^{5} and R^{7} may tomembered heterocyclic ring,

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in still another prefered embodiment,

SUBSTITUTE SHEET (RULE 26)

WO 03/078446

7

PCT/DK03/00176

tionally substituted with one or more substituents selected from the group consisting NR⁵C(=0)R⁸, NR⁵C(=0)OR⁸, NR⁵C(=0)NR⁸R⁷, C(=0)R⁵, C(=NOR⁵)R⁸, C(=0)OR⁵, R³ and R⁴ independently is H, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl opof F, CI, CN, CF3, =0, OR⁶, S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR⁵R⁶, NO₂, NR⁵R⁶,

C(=O)NR5R8, C(=O)NR5OR8 or R8, S

wherein,

gether form a 3-8 membered heterocyclic ring or ${\bf R}^6$ and ${\bf R}^7$ may together form a 3-8 R° , R° , R^{7} and R° independently is H, methyl, ethyl, propyl or butyl and wherein R° and R^{θ} may together form a 3-8 membered heterocyclic ring or R^{δ} and R^{7} may to-

membered heterocyclic ring, 9 in still another prefered embodiment,

cyclopentyl or cyclohexyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF3, =0, OR $^{\!5},\,S(=0)_R^{\!5},\,S(=0)_2R^{\!5},$ R³ and R⁴ independently is methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, $S(=O)_2NR^3R^3$, NO_2 , NR^5R^4 , $NR^5C(=O)R^3$, $NR^5C(=O)NR^3R^7$, C(=0)R⁶, C(=NOR⁵)R⁶, C(=0)OR⁶, C(=0)NR⁵R⁶, C(=0)NR⁵OR⁶ or R⁵,

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Re, Re, R7 and R8 independently is H, cyclopropył, cyclobutyl, cyclopentyl or cyclowherein,

hexyl,

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in still another prefered embodiment,

pholinyl optionally substituted with one or more substituents selected from the group R3 and R4 independently is aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or morconsisting of F, CI, CN, CF₃, =O, OR⁶, S(=O)R⁶, S(=O)₂R⁶, S(=O)₂NR⁶R⁶, NO₂,

NR[‡]R°, NR[‡]C(=0)R°, NR[‡]C(=0)OR°, NR[‡]C(=0)NR°R7, C(=0)R⁵, C(=NOR⁵)R°, C(=0)OR⁵, C(=0)NR⁵R⁶, C(=0)NR⁵OR⁶ or R⁸,

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R⁵, R⁸, R⁷ and R⁸ independently is H, cyclopropył, cyclobutyl, cyclopentyl or cyclo-

hexyl, റ്റ in still another prefered embodiment,

R³ and R⁴ independently is phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, =0, OR 5 , S(=0)R 5 , S(=0)₂N 5 , S(=0)₂NR 5 R 5 ,

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NO2, NR⁵R°, NR˚C(=O)R°, NR˚C(=O)OR°, NR˚C(=O)NR˚R7, C(=O)R˚, $C(=NOR^6)R^6$, $C(=O)OR^5$, $C(=O)NR^6R^6$, $C(=O)NR^5OR^6$ or R^8 ,

Re, R9, R7 and R8 independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclo-

hexyl,

in still another prefered embodiment,

R³ and R⁴ independently is phenyl or naphtyl optionally substituted with one or more S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR⁵R⁵, NO₂, NR⁵R⁵, NR⁵C(=0)R⁶, NR⁵C(=0)OR⁶, substituents selected from the group consisting of F, Cl, CN, CF_{3_i} =0, OR^5 ,

 $\mathsf{NR}^5\mathsf{C}(=\mathsf{O})\mathsf{NR}^6\mathsf{R}^7,\;\;\mathsf{C}(=\mathsf{O})\mathsf{R}^5,\;\mathsf{C}(=\mathsf{NOR}^5)\mathsf{R}^6,\;\mathsf{C}(=\mathsf{O})\mathsf{ON}^5,\;\mathsf{C}(=\mathsf{O})\mathsf{NR}^5\mathsf{OR}^8$

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wherein,

R⁵, R⁶, R⁷ and R⁸ independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclo-

hexyl, 5 in still another prefered embodiment,

R³ and R⁴ independently is thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally CI, CN, CF3, =0, OR¢, S(=0)R[¢], S(=0)₂R[¢], S(=0)₂NR⁵R[¢], NO₂, NR⁵R[¢], NR⁵C(=0)R[¢] NR⁵C(=0)OR°, NR⁵C(=0)NR°R7, C(=0)R⁵, C(=NOR⁵)R°, C(=0)OR⁵, C(=0)NR°R°, substituted with one or more substituents selected from the group consisting of F, ឧ

C(=O)NR⁵OR⁸ or R⁸,

R⁵, R⁶, R⁷ and R⁸ independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclowherein,

hexyl, 22 in still another prefered embodiment,

cyclopentyl or cyclohexyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF3, =0. OR5 , $S(=O)R^5$, $S(=O)_2R^5$, R³ and R⁴ independently is methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, S(=0)2NR5R°, NO2, NR5R°, NR5C(=0)R°, NR°C(=0)OR°, NR°C(=0)NR®R7, C(=0)R⁶, C(=NOR⁵)R⁶, C(=0)OR⁵, C(=0)NR⁵CR⁶ or R⁸,

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R^s, R^s, R² and R⁸ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, qui

nolinyl or isoquinolinyl,

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SUBSTITUTE SHEET (RULE 26)

WO 03/078446

33

PCT/DK03/00176

in still another prefered embodiment,

pholinyl optionally substituted with one or more substituents selected from the group R³ and R⁴ independently is aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or mor-

NR°R°, NR°C(=0)R°, NR°C(=0)OR°, NR°C(=0)NR°R7, C(=0)R°, C(=NOR°)R°, consisting of F, Cl, CN, CF_{3,} =O, OR⁵, S(=O)R⁵, S(=O)₂R⁵, S(=O)₂NR⁵R⁸, NO₂. $C(=O)OR^5$, $C(=O)NR^5R^9$, $C(=O)NR^5OR^9$ or R^8 , ß

R⁵, R⁶, R⁷ and R⁸ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, qui-

nolinyl or isoquinolinyl,

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in still another prefered embodiment,

R³ and R⁴ independently is phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the

group consisting of F, Cl, CN, CF₃, =O, OR⁵, S(=O)R⁵, S(=O)₂NR⁵R⁶, NO2, NR⁵R⁶, NR⁵C(=0)R⁶, NR⁵C(=0)NR⁵R, C(=0)R⁵, $C(=NOR^5)R^6$, $C(=O)OR^5$, $C(=O)NR^5R^6$, $C(=O)NR^5OR^6$ or R^5 , रु

R⁵, R⁶, R⁷ and R⁸ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, qui-

nolinyl or isoquinolinyl,

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in still another prefered embodiment,

R³ and R⁴ independently is phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF₃, =O, OR⁵,

NR⁵C(=O)NR⁶R⁷, C(=O)R⁵, C(=NOR⁵)R⁶, C(=O)OR⁵, C(=O)NR⁵R⁶, C(=O)NR⁵OR⁶ S(=0)R[¢], S(=0)₂R[¢], S(=0)₂NR[¢]R[¢], NO₂, NR[¢]R[¢], NR[¢]C(=0)R[¢], NR[¢]C(=0)OR[¢], م ہے 22

wherein,

R⁵, R⁶, R⁷ and R⁸ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, qui-

nolinyl or isoquinolinyl, ဓ in still another prefered embodiment,

R³ and R⁴ independently is thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F,

CI, CN, CF3, =0, OR⁶, S(=0)R⁵, S(=0)₂R⁵, S(=0)₂NR⁵R⁶, NO₂, NR⁵R⁶, NR⁵C(=0)R⁶, 33

24

NR⁶C(=0)OR⁸, NR⁶C(=0)NR⁸R⁷, C(=0)R⁵, C(=NOR⁶)R⁶, C(=0)OR⁵, C(=0)NR⁵R⁶,

C(=0)NR5OR8 or R8,

wherein,

R^e, R^o, R⁷ and R⁸ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, qui-

nolinyl or isoquinolinyl,

in still another prefered embodiment,

 R^3 and R^4 independently is H, $C_1\text{-}C_6$ alkyl, $C_3\text{-}C_7$ cycloalkyl, $C_3\text{-}C_7$ cycloheteroalkyl,

aryl or heteroaryl

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in still another prefered embodiment,

R3 and R4 independently is H,

in still another prefered embodiment,

R3 and R4 independently is C1-C8 alkyl, C3-C7 cycloalkyl or C3-C7 cycloheteroalkyl 5

in still another prefered embodiment,

R3 and R4 independently is methyl, ethyl, propyl or butyl

in still another prefered embodiment ೫ R3 and R4 independently is cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl

in still another prefered embodiment

R3 and R4 independently is aziridinyl, pyrrolidinyl, piperidinyl or morpholinyl

in still another prefered embodiment,

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R3 and R4 independently is aryl or heteroaryl

in still another prefered embodiment,

R3 and R4 independently is phenyl or naphthyl ဓ

R3 and R4 independently is thienyl, furyl, pyridyl, quinolinyl or isoquinolyl

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in still another prefered embodiment,

WO 03/078446

23

PCT/DK03/00176

Experimental section

General Procedure 1: Preparation of Carrier-Functional entity reagents:

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hours. The mixture is added to 100 mL ice and the precipitate collected by filtration. he 4-halobenzoic acid (25 mmol) is added to a ice cooled solution of chloro sulfonic acid (140 mmol). The mixture is slowly heated to reflux and left at reflux for 2-3

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cooled mixture of NaOEt (10 mL, 2M). The mixture is stirred o/n at rt. Acetic acid (40 mmol) is added and the mixture is evaporated in vacuo. Water (10 mL) is added and pH adjusted to pH = 2 (using 1M HCI). The product is extracted with DCM (2 x25 The filtrate is washed with water (2 \times 50 mL) and the dried in vacuo affording the mL), dried over Na₂SO₄ and evaporated in vacuo affording the desired products. benzoic acid derivate (5 mmol) is dissolved in EtOH (5 mL) and added to a ice corresponding sulfonoyl chloride in 60-80% yield. The 3-chlorosulfonyl-4-halo-

Example 1 (General procedure (1))

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3-Ethoxysulfonyl-4-fluorobenzoic acid

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H-NMR (DMSO-d₆): 5 8.49 (d, 1H), 7.85 (dd, 1H), 7.5 (d, 1H), 4.32 (q, 2H), 1.32 (t,

4-chloro-3-Ethoxysulfonylbenzoic acid

Example 2 (General procedure (1))

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H-NMR (DMSO-46): 5 8.49 (d, 1H), 7.85 (dd, 1H), 7.5 (d, 1H), 4.32 (q, 2H), 1.32 (t,

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PCT/DK03/00176

58

Example 3

4-Methylsulfanyl benzoic acid (0.5g, 2.97 mmol, commercially available from Aldrich, cat no. 145521) was added to methyl p-toluene solfunate (0.61g, 3.27 mmol). The mixture was heated to 140 °C for 1 hour in a sealed vessel. After cooling to rt the mixture was trituated with diethyl ether. Filtration and drying *in vacuo* yielded 844 mg (80%) of the desired product (>95% pure by ¹H nmr).

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¹H nmr (DMSO-d6): 8.20-8.10 (m, 4H), 7.45 (d, 2H), 7.08 (d, 2H), 3.29 (s, 6H), 2.30 (s, 3H).

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General Procedure 2: Solid phase preparation of Carrier-Functional entity reagents

for alkylation building blocks:

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Ps = Polystyrene resin. Alternatively other acid labile linkers may be employed.

Step 1:

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A polystyrene resin with a wang linker (4-hydroxymethylphenol linker) (50 mg \sim 50 unol), a bl-functional carrier (200 umol, 4 equiv) in a solvent such as THF, DCM.

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WO 03/078446

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PCT/DK03/00176

DCE, DMF, NMP or a mixture thereof (500 uL) and a base such as TEA, DIEA, pyridine (400 umol, 8 equiv), optionally in the presence of DMAP (100 umol), are allowed to react at temperatures between -20 °C and 60 °C, preferably between 0 °C and 25 °C, for 1-24 h, preferably 1-4 h. The resin is washed with the solvent composition used during the reaction (5x1 mL) and used in the following step.

Step 2:

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A functional entity precursor carrying a hydroxy group in the position of the intended attachment to the C-F-connecting group (200 umol, 4 equiv) in a solvent such as THF, DCM, DCE, DMF, NMP or a mixture thereof (500 uL) and a base such as TEA, DIEA, pyridine (400 umol, 8 equiv), optionally in the presence of DMAP, are added to the resin bound carrier isolated in step 1 and allowed to react at temperatures between 0 °C and 100 °C, preferably between 25 °C and 80 °C, for 2-48 h, preferably 4-16 h. The resin is washed with the solvent composition used during the reaction (5x1 mL).

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tep 3:

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The desired Carrier-Functional entity reagent is cleaved from the resin obtained in step 2 by treatment with an acid like TFA, HF or HCl in a solvent such as THF, DCM, DCE or a mixture thereof (1 mL) at temperatures between -20 °C and 60 °C, preferably between 0 °C and 25 °C, for 1-4 h, preferably 1-2 h. Upon filtration, the resin is washed with the solvent composition used during cleavage (2x1 mL) and the combined filtrates are evaporated in vacuo. The isolated product may be purified by chromatography.

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Assembly of building blocks

The Carrier-Functional entity reagent may be bound to the Spacer by several different reactions as illustrated below.

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Formation of an amide bond between a carboxylic acid of the Carrier and an amine group of a Spacer

General Procedure 3: Preparation of building blocks by loading a Carrier-Functional entity reagent onto a nucleotide derivative comprising an amino group:

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15 µL of a 150 mM building block solution of FE¹-Carrier-COOH is mixed with 15 µL of a 150 mM solution of EDC and 15 µL of a 150 mM solution of N-hydroxysuccinimide (NHS) using solvents like DMF, DMSO, water, acetonitril, THF, DCM, methanol, ethanol or a mixture thereof. The mixture is left for 15 min at 25°C. 45 µL of an aminooligo (10 nmol) in 100 mM buffer at a pH between 5 and 10, preferably 6.0-7.5 is added and the reaction mixture is left for 2 hours at 25°C. Excess building block and organic by-products were removed by extraction with EtOAc (400 µL). Remaining EtOAc is evaporated *in vacuo* using a speedvac. The building block is purified following elution through a BioRad micro-spin chromatography column, and analyzed by electron spray mass spectrometry (ES-MS).

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Example 4 (General procedure ())

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Where Oligo is 5' XCG ATG GAT GCT CCA GGT CGC 3', X = 5' amino C6 (Glen catalogue# 10-1906-90), Expected molecular weight: 6313.22

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WO 03/078446

53

PCT/DK03/00176

MS (calc.) = 6543,43; MS (found) = 6513,68*

 Observed molecular weight of the chaved suitonic ester. 6513.88 Expected molecular weight of the cleaved ester.
 6514.37 The quantitative loss of the ethyl group is probably due to the presence of piperidine during the recording of the LC-MS data.

General Procedure 4: Loading of a carrier coupled functional entity onto an amino

oligo:

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25 µl 100 mM carrier coupled functional entity dissolved in DMF (dimethyl formamide) was mixed with 25 µl 100 mM EDC (1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride) in DMF for 30 minutes at 25° C. The mixture was added to 50 µl amino oligo in H₂O with 100 mM HEPES (2-(4-(2-hydroxy-ethyl))-piperazin-1-yl]-ethanesulfonic acid) pH 7.5 and the reaction was allowed to proceed for 20 minutes at 25° C. Unreacted carrier coupled functional entity was removed by extraction with 500 µl EtOAc (ethyl acetate), and the oligo was purified by gel filtration through a microspin column equilibrated with 100 mM MES (2-(N-morpholino) ethanesulfonic

Oligonucleotide used:

acid) pH 6.0.

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Oligo A: 5'-YACGATGGATGCTCCAGGTCGC

Y = Amino modifier C6 (Glen# 10-1906)

Example 5 (General procedure 4)
Carrier - Functional Entity: (4-Carboxy-phenyl)-dimethyl-sulfonium

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Mass: 6789.21 (observed using ES-MS), 6790.65 (calculated)

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General Procedure 5: Preparation of arylation building blocks:

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Funtional Entity-OH is a phenol, n is an integer between 3 and 6.

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functional entity in excess (1.05-1.8 mmol) in DMF, DMSO, acetonitril, THF or a mix-To a solution of the bis-sulfonylchloride (Ward, R.B.; J.Org.Chem.; 30; 1965; 3009-3011; Qiu, Weiming; Burton, Donald J.; J.Fluorine Chem.; 60; 1; 1993; 93-100) (3 umol) in DMF, DMSO, acetonitril, THF or a mixture thereof (150 uL) is a phenolic preferably at 0-50 °C in the presence of a base such as TEA, DIEA, pyridine, Nature thereof (150 u.L.) added slowly at temperatures between -20 °C and 100 °C HCO3 or K2CO3.

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ing EtOAc is evaporated in vacuo using a speedvac. The building aminooligo is puri-The reaction mixture from step 1 is added to a solution of an aminooligo (10 nmol) in and organic by-products were removed by extraction with EtOAc (400 µL). Remainence of NHS. The reaction mixture is left for 2 hours at 25°C. Excess building block 100 mM buffer at a pH between 5 and 10, preferably 6.0-7.5 optionally in the presfied following elution through a BioRad micro-spin chromatography column, and analyzed by electron spray mass spectrometry (ES-MS).

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WO 03/078446

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PCT/DK03/00176

Use of building blocks

General Procedure 6: Alkylation of oligonucleotide derivatives containing a nucleophilic recipient group using a building block of the invention.

-recipient reactive group

a nucleophilic recipient group. Reaction proceeds at temperatures between 0 °C and final concentration with one equivalent of a complementary building block displaying pH buffered to 4-10, preferably 6-8. Organic by-products are removed by extraction with EtOAc, followed by evaporation of residual organic solvent for 10 min in vacuo. An oligonucleotide building block carrying functional entity FE¹ is combined at 2 µM Pd catalyst is removed and oligonucleotides are isolated by eluting sample through DMF, DMSO, water, acetonitril, THF, DCM, methanol, ethanol or a mixture thereof, 100 °C preferably between 15 °C-50 °C for 1-48 hours, preferably 10-20 hours in a BioRad micro-spin chromatography column. Coupling efficiency is quantified by ES-MS analysis. 5 9

General procedure 7: Transfer of functional entity from a carrier oligo to recipient

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microspin column equilibrated with H2O and transfer of the functional entity was verified by electron spray mass spectrometry (ES-MS). Transfer efficiency is expressed scaffold oligo B (200 pmol) in 50 µl 100 mM MES, pH 6. The mixture was incubated transferred functional entities to total abundance of scaffold oligos (with and without n percent and were calculated by dividing the abundance of scaffold oligo carrying overnight at 25 °C. Subsequently, the mixture was purified by gel filtration using a A carrier coupled functional entity oligo (Example 1) (250 pmol) was added to a transferred functional entities).

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Example 6 (General procedure 7) ဓ္က

PCT/DK03/00176

Mass ("X"): 6583.97 (observed), 6583.31 (calculated). Abundance: 65.79 (arbitrary

- Mass ("Y"): 6599.73 (observed), 6597.34 (calculated). Abundance: 29.23 (arbitrary
- Mass ("Z"): 6789.36 (observed), 6790.65 (calculated)
- Transfer efficiency calculated as: 29.23 / (29.23 + 65.79) = 0.3076 \sim 31 % 6

WO 03/078446

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PCT/DK03/00176

General Procedure 8: Arylation of oligonucleotide derivatives containing a nucleophilic recipient group using a building block of the invention:

X=recipient reactive group

final concentration with one equivalent of a complementary building block displaying Organic by-products are removed by extraction with EtOAc, followed by evaporation of residual organic solvent for 10 min in vacuo. Pd catalyst is removed and oligonu-An oligonucleotide building block carrying functional entity FE¹ is combined at 2 µM cleotides are isolated by eluting sample through a BioRad micro-spin chromatograceeds at temperatures between 0 °C and 100 °C preferably between 15 °C-50 °C DCM, methanol, ethanol or a mixture thereof, pH buffered to 4-10, preferably 6-8. a nucleophilic recipient group. In the presence of a Pd catalyst, the reaction profor 1-48 hours, preferably 10-20 hours in DMF, DMSO, water, acetonitrile, THF, phy column. Coupling efficiency is quantified by ES-MS analysis.

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monomer building blocks with a thio-succinimid S-C-connecting group and use of General Procedure 9: General route to the formation of alkylating/vinylating

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 $R^1 = H$, Me, Et, iPr, CI, NO₂ $R^2 = H$, Me, Et, iPr, CI, NO₂

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R1 and R2 may be used to tune the reactivity of the sulphate to allow appropriate reactivity. Chloro and nitro substitution will increase reactivity. Alkyl groups will decrease reactivity. Ortho substituents to the sulphate will due to steric reasons direct incoming nucleophiles to attack the R-group selectively and avoid attack on sulphur.

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3-Aminophenol (6) is treated with maleic anhydride, followed by treatment with an acid e.g. H₂SO₄ or P₂O₅ and heat to yield the maleimide (7). The ring closure to the maleimide may also be achieved when an acid stable O-protection group is used by treatment with or Ac2O with or without heating, followed by O-deprotection. Alternatively reflux in Ac₂O, followed by O-deacetylation in hot water/dioxane to yield (7).

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the quench with the appropriate alcohol, in this case MeOH, whereby (9) will be formed. The organic building block (9) may be connected to an oligo nucleotide, as Further treatment of (7) with SO₂Cl₂ with or without triethylamine or potassium carbonate in dichloromethane or a higher boiling solvent will yield the intermediate (8), which may be isolated or directly further transformed into the aryl alkyl sulphate by

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A thiol carrying oligonucleotide in buffer 50 mM MOPS or hepes or phosphate pH 7.5 is treated with a 1-100 mM solution and preferably 7.5 mM solution of the organic building block (9) in DMSO or alternatively DMF, such that the DMSO/DMF concentration is 5-50%, and preferably 10%. The mixture is left for 1-16 h and prefsrably 2-4 h at 25 \circ C. To give the alkylating in this case methylating monomer building block (10).

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The reaction of the alkylating monomer building block (10) with an amine carrying monomer building block may be conducted as follows:

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(20 μL of a 100 mM hepes and 1 M NaCl solution, pH=7.5) and water (39 uL). The oligonucleotides are annealed to the template by heating to 50 °C and cooled (2 °C/ second) to 30 °C. The mixture is then left o/n at a fluctuating temperature (10 °C for The coding oligonucleotide (1 nmol) is mixed with a thio oligonucleotide loaded with a building block (1 nmol) (10) and an amino-oligonucleotide (1 nmol) in hepes-buffer 1 second then 35 °C for 1 second), to yield the template bound methylamine (11).

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the chlorosulphonate (8 above) with an alcohol, the intermediate chlorosulphate is scribed above for an alkylating monomer building block. Although instead of reacting isolated and treated with an enolate or O-trialkylsilylenolate with or without the pres-A vinylating monomer building block may be prepared and used similarily as deence of fluoride. E.g.

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Formation of the vinylating monomer building block (13):

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The thiol carrying oligonucleotide in buffer 50 mM MOPS or hepes or phosphate pH 7.5 is treated with a 1-100 mM solution and preferably 7.5 mM solution of the organic building block (12) in DMSO or alternatively DMF, such that the DMSO/DMF concentration is 5-50%, and preferably 10%. The mixture is left for 1-16 h and preferably 2-4 h at 25 °C. To give the vinylating monomer building block (13).

The sulfonylenolate (13) may be used to react with amine carrying monomer building block to give an enamine (14a and/or 14b) or e.g. react with an carbanion to yield (15a and/or 15b). E.g.

WO 03/078446

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The reaction of the vinylating monomer building block (13) and an amine or nitroal-kyl carrying monomer building block may be conducted as follows:

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The coding oligonucleotide (1 nmol) is mixed with a oligonucleotide building block (1 nmol) (13) and an amino-oligonucleotide (1 nmol) or nitroalkyl-oligonucleotide (1 nmol) in 0.1 M TAPS, phosphate or hepes-buffer and 300 mM NaCl solution, pH=7.5-8.5 and preferably pH=8.5. The oligonucleotides are annealed to the template by heating to 50 °C and cooled (2 °C/ second) to 30 °C. The mixture is then left o/n at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second), to yield template bound (14a/b or 15a/b).

9

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Abbreviations

DCC	N,N'-Dicyclohexylcarbodiimide
DhbtOH	3,4-dihydro-3-hydroxy-4-oxo-1,2,3-benzotriazine
DIC	Diisopropylcarbodiimide
DIEA	Diethylisopropylamin
DMAP	4-Dimethylaminopyridine

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Claims

Complementing Element - Linker - Carrier - C-F-connecting group - Func-

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2-(1H-Benzotriazole-1-yl)-1,1,3,3-tetramethyluronium hexafluoro-

N-Hydroxy-7-azabenzotriazole

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phosphate

HBTU

N-Hydroxybenzotriazole

Frifluoromethylsulfonate

N-hydroxysuccinimid Locked Nucleic Acid

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2-(1H-7-Azabenzotriazole-1-yl)-1,1,3,3-tetramethyluronium

HATU

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hexafluorophosphate

1-Ethyl-3-(3'-dimethylaminopropyl)carbodiimide·HCl

Deoxyribosenucleic Acid

capable of transferring a Functional entity precursor to a recipient reactive group,

Complementing Element is a group identifying the Functional entity precursor,

entity precursor to be transferred from the complementing element and the S-C-Linker is a chemical moiety comprising a spacer and a S-C-connecting

9

kynylene, or -(CF2) $\!\!\!\!^{-}$ substituted with 0-3 R¹ wherein m is an integer between 1 and

-C(Halogen)₃, -C(O)R², -C(O)NHR², C(O)NR²₂, -NC(O)R², -S(O)₂NHR², -S(O)₂NR²₂, -S(O)₂R², _P(O)₂-R², _P(O)- R², _S(O)- R², P(O)-OR², _S(O)-OR², -N¹R²₃, wherein R¹ are independently selected from -H, -OR², -NR²₂, -Halogen, -NO₂, -CN,

-O-SO₂-O-, -C(O)-O-, -S*(R³)-, -C-U-C(V)-O-, -P*(W)z-O-, -P(W)-O- where U is C-F-connecting group is chosen from the group consisting of -SO₂-O-, -C(R²)_{2~}, -NR²- or –O-; V is =O or =NR² and W is -OR² or –N(R²)₂

eroaryl or aryl optionally substituted with one or more substituents belonging to the Functional entity precursor is -C(H)(R³)-R⁴ or functional entity precursor is hetgroup comprising R3 and R4.

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Boc anhydride, di-tert-butyl dicarbonate

(Boc)₂O

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TBAF

SPDP

Thin layer chromatography

5-iodo-deoxyriboseuracil

Succinimidyl-propyl-2-dithiopyridyl

Tetrabutylammonium fluoride

cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substitu-Sn(OR³)(OR³)R7, BR⁵R⁵, B(OR⁵)R⁴, B(OR⁵)(ORª), halogen, CN, CNO, C(halogen)₃, S(=O)2NR5R⁶, NO2, N3, NR⁵R⁶, N⁺R⁶R⁷, NR⁵OR⁶, NR⁵NR⁶R⁷, NR⁵C(=O)R⁶, OR⁵, OC(=0)R⁵, OC(=0)OR⁵, OC(=0)NR⁵R³, SR⁵, S(=0)R⁵, S(=0)₂R⁵, ents selected from the group consisting of SnR⁵R⁶R⁷, Sn(OR⁵)R⁶R⁷,

1. A building block of the general formula

tional entity precursor

group, wherein the spacer is a valence bond or a group distancing the Functional connecting group connects the spacer with the Carrier

Carrier is arylene, heteroarylene, C.-C. alkylene, C.-C. alkenylene, C.-C. al-

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Benzotriazole-1-yl-oxy-tris-pyrrolidino-phosphonium hexafluoro-

Peptide Nucleic Acid **Foluenesulfonate**

2-(1 H-Benzotriazole-1-yl)-1,1,3,3-tetramethyluronium tetra-

Ribonucleic acid

Bromo-tris-pyrrolidino-phosphonium hexafluorophosphate

phosphate

PyBoP

₽N ₩

OTs

OTf

PyBroP

TBTU

RN ₩

Reverse Phase High Performance Liquid Chromatography

Friethylamine fluoroborate

Tert-Butyldimethylsilylchloride

TBDMS-CI

5-lodo-dU

RP-HPLC

IEA

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R2 is H, C1-C6 alkyl, C2-C6 alkenyl, C2-C6 alkynyl, or aryl,

Wherein R³ and R⁴ independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl,

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NR⁶C(=0)OR⁶, NR⁵C(=0)NR⁸R⁷, NC, P(=0)(OR⁶)OR⁶, P⁺R⁵R⁵R⁷, C(=0)R⁵,

A

C(=NR³)R°, C(=NOR⁵)R°, C(=NNR⁵R°), C(=O)OR°, C(=O)NR⁵R°, C(=O)NR⁵OR° C(=O)NR⁶NR⁶R⁷, C(=NR⁵)NR⁶R⁷, C(=NOR⁵)NR⁶R⁷ or R⁸,

2

cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents selected from the group consisting of halogen, CN, CNO, C(halogen)3, =O, R^{ς} , R^{ε} , and R^{γ} independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl, S(=0)¿NR®R°, NO2, N3, NR®R°, N*R®R°R¹º, NR°OR°, NR°NR®R', NR°C(=0)R°, NR®C(=0)OR®, NR®C(=0)NR®R®, NC, P(=0)(OR®)OR®, P*R®R7, C(=0)R®, OR", OC(=0)R", OC(=0)OR", OC(=0)NR"R", SR", S(=0)R", S(=0)2R",

gether form a 3-8 membered heterocyclic ring or $\ensuremath{R^{5}}$ and $\ensuremath{R^{7}}$ may together form a 3-8 membered heterocyclic ring or $\ensuremath{R^{a}}$ and $\ensuremath{R^{7}}$ may together form a 3-8 membered het-C(=NR5)NR6R7, C(=NOR5)NR6R7 or C(=O)NR6NR9R10, wherein R5 and R6 may to-C(=NR³)R°, C(=NOR³)R°, C(=NNR³R³), C(=O)OR°, C(=O)NR³R°, C(=O)NR³OR° erocyclic ring,

2

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cycloheteroalkyl, aryl or heteroaryl and wherein R^{θ} and R^{θ} may together form a 3-8 $\,$ membered heterocyclic ring or R^{9} and R^{10} may together form a 3-8 membered heterocyclic ring or R^{θ} and R^{10} may together form a 3-8 membered heterocyclic ring. R⁸, R⁹, and R¹⁰ independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl,

-C(H)(R11)-R11 or functional entity precursor is heteroaryl or aryl substituted with 0-3 2. A compound according to claim 1 wherein, Functional entity precursor is R11, 0-3 R13 and 0-3 R15, wherein

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 R^{11} and R^{11} are independently H, or selected among the group consisting of a $\mathsf{C}_1\text{-}\mathsf{C}_6$ heteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 $m R^{12}$, 0-3 $m R^{13}$ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₆ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cyclo-

22

or R¹¹ and R¹¹ are C₁-C₃ alkylene-NR¹², C₁-C₃ alkylene-NR¹²C(O)R¹⁹, C₁-C₃ alkylene-NR¹²C(O)OR¹⁶, C₁-C₂ alkylene-O-NR¹², C₁-C₂ alkylene-O-NR¹²C(O)R¹⁶, C₁-C₂ alkylene-O-NR¹²C(O)OR¹⁶ substituted with 0-3 R¹⁵,

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where R¹² is H or selected independently among the group consisting of C₁-C₈ alkyl, C2-Ce alkenyl, C2-Ce alkynyl, C3-C, cycloalkyl, C3-C, cycloheteroalkyl, aryl, heteroaryl, said group being substituted with 0-3 \mbox{R}^{13} and 0-3 $\mbox{R}^{15},$

-NHNHR1", -C(O)R1", -SnR1"3, -B(OR17)2, -P(O)(OR17)2 or the group consisting of R13 is selected independently from -N3, -CNO, -C(NOH)NH2, -NHOH,

35

WO 03/078446

PCT/DK03/00176

 $C_{z}\dot{-}C_{g}$ alkenyl, $C_{z}C_{g}$ alkynyl, $C_{d}\dot{-}C_{g}$ alkadienyl said group being substituted with 0-2

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where R14 is independently selected from -NO2, -C(0)OR17, -COR17, -CN, -OSiR¹⁷3, -OR¹⁷ and -NR¹⁷2;

-NR"7-C(O)OR16, -SR17, -S(O)R17, -S(O)2R17, -COOR17, -C(O)NR172 and -S(O)2NR172, R¹⁶ is H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, aryl or C₁-C₆ alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -R¹⁵ is =O, -F, -CI, -Br, -I, -CN, -NO₂, -OR¹⁷, -NR¹⁷, -NR¹⁷-C(O)R¹⁶, NO2, -R2, -OR2, -SiR3;

R¹⁷ is selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, aryl, C₁-C₆ al-9

Kylene-aryl, Cot or Cha, G is H or Ct-Ce alkyl and n is 1,2,3 or 4.

-C(H)(R11)-R11 or functional entity precursor is heteroaryl or aryl substituted with 0-3 3. A compound according to claim 2 wherein, Functional entity precursor is

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R11 and R11, are independently H, or selected among the group consisting of a C1-Ce heteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 $m R^{12}$, 0-3 $m R^{13}$ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₆ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cyclo-R11, 0-3 R13 and 0-3 R15, wherein and 0-3 R¹⁵,

or R¹¹ and R¹¹, are C₁-C₃ alkylene-NR¹², C₁-C₃ alkylene-NR¹²C(O)R¹⁹, C₁-C₃ alkylene-NR12C(O)OR19, C;-C2 alkylene-O-NR12, C;-C2 alkylene-O-NR12C(O)R16, C₁-C₂ alkylene-O-NR¹²C(O)OR¹⁸ substituted with 0-3 R¹⁵,

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where R12 is H or selected independently among the group consisting of C1-C8 alkyl, C₂-C₈ alkenyl, C₂-C₈ alkynyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl, heteroaryl, said group being substituted with 0-3 $\ensuremath{\text{R}}^{13}$ and 0-3 $\ensuremath{\text{R}}^{15}$

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C₂-C₈ alkenyi, C₂-C₈ alkynyi, C₄-C₈ alkadienyi said group being substituted with 0-2 NHNHR17, -C(O)R17, -SnR17s, -B(OR17)2, -P(O)(OR17)2 or the group consisting of R13 is selected independently from -N3, -CNO, -C(NOH)NH2, -NHOH,

where R^{14} is independently selected from $-NO_2,\, -C(O)OR^{17},\, -CON,$ -OSIR¹⁷3, -OR¹⁷ and -NR¹⁷2;

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 $-NR^{17}$ -C(O)OR¹⁸, $-SR^{17}$, $-S(O)_2R^{17}$, $-COOR^{17}$, $-C(O)NR^{12}$ and $-S(O)_2NR^{12}$. R¹⁶ is =0, -F, -CI, -Br, -I, -CN, -NO₂, -OR¹⁷, -NR¹⁷, -NR¹⁷-C(O)R¹⁶,

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alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -R¹⁸ is H, C₁-C₈ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, aryl or C₁-C₆ NO2, -R2, -OR2, -SIR23;

wherein R¹⁷ is selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, aryl,

C₁-C₆ alkylene-aryl.

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4. A compound according to claim 1 wherein, Functional entity precursor is -C(H)(R11)-R111 wherein

 R^{11} and R^{11} are or $C_1\text{-}C_3$ alkylene-NR $^{12}\!,$ $C_1\text{-}C_3$ alkylene-NR $^{12}\!C(O)R^{16}\!,$ $C_1\text{-}C_3$ alkylene-NR¹²C(O)OR¹6, C₁-C₂ alkylene-O-NR¹2, C₁-C₂ alkylene-O-NR¹²C(O)R¹6

C₁-C₂ alkylene-O-NR¹²C(O)OR¹⁸ substituted with 0-3 R¹⁵

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5. A compound according to claim 1 wherein, Functional entity precursor is -C(H)(R11)-R111 wherein

 R^{11} and R^{11} are independently H, or selected among the group consisting of a $C_1\text{-}C_6$ heteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 $m R^{12}$, 0-3 $m R^{13}$ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₆ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloand 0-3 R¹⁵,

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6. A compound according to claim 2 wherein, Functional entity precursor is -C(H)(R11)-R111 wherein 2

R11 and R11, are independently H, or selected among the group consisting of a C1-Ce alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R¹² and 0-3 R¹⁵,

where R12 is H or selected independently among the group consisting of C1-C6 alkyi, C2-Ce alkenyi, C2-Ce alkynyi, C3-C7 cycloalkyi, C3-C7 cycloheteroalkyi, aryi,

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R¹⁵ is =0, -F, -CI, -Br, -I, -CN, -NO₂, -OR¹⁷, -NR¹⁷, -NR¹⁷-C(0)R¹⁶,

-NR 17 -C(O)OR 16 , -SR 17 , -S(O)R 17 , -S(O)₂R 17 , -COOR 17 , -C(O)NR 12 and -S(O)₂NR 7 ₂, R¹⁷ is selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₁-C₆ al-ဓ

7. A compound according to claim 1 wherein, Functional entity precursor is heteroaryl or aryl substituted with 0-3 R11, 0-3 R13 and 0-3

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WO 03/078446

PCT/DK03/00176

-P*(W)z-O., and -P(W)-O- where U is -C(R²)z-, -NR²- or -O-; V is =O or =NR² and W A compound according to claim 2 wherein C-F-connecting group is chosen from the group consisting of -SO₂-O-, -O-SO₂-O-, -C(O)-O-, -S*(R*1)-, -C-U-C(V)-O-, is -OR2 or -N(R2)2

9. A compound according to claim 2 wherein C-F-connecting group is -S*(R¹¹)-,

0. A compound according to claims 1 - 2 wherein C-F-connecting group is chosen from the group consisting of -SO₂-O-, -O-SO₂-O-, -C(O)-O-, -S*(R¹⁷)-, -C-U-

 $C(V)-O_+$, -P*(W)₂-O-, and -P(W)-O- where U is -C(R²)₂-, -NR²- or -O-; V is =O or =NR 2 and W is -OR 2 or –N(R $^2\rangle_2$, wherein R 17 is selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, aryl, C₁-C₆ alkylene-aryl. 9

11. A compound according to claims 1 - 2 wherein C-F-connecting group is chosen from the group consisting of -SO₂-O-, and -S*(R¹7)-; wherein R¹7 is selected ndependently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, aryl, C₁-C₆ alkylene-aryl.

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 A compound according to claim 1 wherein, Spacer is a valence bond, C₁-C₆ alkylene-A-, C2-C6 alkenylene-A-, C2-C6 alkynylene-A-, or

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said spacer optionally being connected through A to a linker selected from

$$-(CH_2)_n-B-$$
, and

--(CH₂)_n-S-S-(CH₂)_m-B--

where A is a valence bond, -C(O)NR¹⁷-, -NR¹⁷-, -O-, -S-, or -C(O)-O-; B is a valence bond, -O., -S., -NR17 or -C(O)NR17 and connects to S-C-connecting group; and n and m independently are integers ranging from 1 to 10; and \mathbb{R}^{17} is selected ndependently from H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, aryl, or C₁-C₈ alkylene-aryl

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13. A compound according to claim 1 wherein, Spacer is a valence bond, C₁-C₆ alkylene-A-, C2-C6 alkenylene-A-, C2-C6 alkynylene-A-, or

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PCT/DK03/00176

said spacer optionally being connected through A to a linker selected from

$$-(CH_2)_n-B-$$
, O n and

where A is a valence bond, —C(O)NR¹²-, -S-, or -C(O)-O-; B is -O-, -S-, -NR³²-, or -C(O)NR³²- and connects to S-C-connecting group; and n and m independently are integers ranging from 1 to 6; and R³ is selected independently from H, C₁-C₀ alkyl, C₃-C, cycloalkyl, aryl, or C₁-C₀ alkylene-aryl

14. A compound according to claim 1-2 wherein, S-C-connecting group is a va-

10 lence bond, –NH-C(=O)-, –NH-SO₂-, -S-S-,

15. A compound according to claim 2 wherein, the carrier is selected from the group consisting of anylene, heteroarylene or $-(CF_2)_{nr}$ substituted with 0-3 R¹ wherein m is an integer between 1 and 10, and C-F-connecting group is $-SO_z-O_-$, and the functional entity precursor is $-C(H)(R^{11})-R^{11}$.

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20 16. A compound according to claim 2 wherein, the carrier is -(CF₂)_m- wherein m is an integer between 1 and 10, the C-F-connecting group is -SO₂-O-; and the functional entity precursor is anyl or heteroaryl substituted with 0-3 R¹¹, 0-3 R¹³ and 0-3 R¹⁵

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WO 03/078446

PCT/DK03/00176

 A compound according to claims 1-16 wherein Complementing element is a nucleic acid.

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18. A compound according to claims 1-16 where Complementing element is a sequence of nucleotides selected from the group of DNA, RNA, LNA PNA, or morpholino derivatives.

19. A library of compounds according to claim 1, wherein each different member of the library comprises a complementing element having a unique sequence of nucleotides, which identifies the functional entity.

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20. A method for transferring a functional entity precursor to a recipient reactive group, comprising the steps of

providing one or more building blocks according to claims 1 to 18,

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contacting the one or more building blocks with a corresponding encoding element associated with a recipient reactive group under conditions which allow for a recognition between the one or more complementing elements and the encoding elements, said contacting being performed prior to, simultaneously with, or subsequent to a transfer of the functional entity precursor to the recipient reactive group.

21. The method according to claim 20, wherein the encoding element comprises one or more encoding sequences comprised of 1 to 50 nucleotides and the one or more complementing elements comprises a sequence of nucleotides complementary to one or more of the encoding sequences.

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22. The method of claims 20 or 21, wherein the recipient reactive group is a nucleo-philic S- or N- atom, which may be part of a chemical scaffold, and the activating catalyst is contains palladium.

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Figure 1. Two setups for Functional Entity Transfer

-Linker-Carrier-F.E. Prec. 1-F.E. Prec. 2 -Linker-Carrier Functional Entity Precursor Transfer Complementing element Complementing element Coding Element Coding Element

Complementing element F.E. Prec. 1 F.E. Prec. 2 Carrier Complementing element Linker

Carrier-Linker Functional Entity Precursor Transfer F.E. Prec. 1-F.E. Prec. 2 Complementing element Linker Coding Element

Coding Element

WO 03/078446

PCT/DK03/00176

Figure 2. Examples of specific base pairing

Natural Base Pairs

Synthetic Base Pairs

Synthetic purine's base pairing with U/T or C

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Figure 3. Example of non-specific base-pairing

I = Inosine

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Figure 4. Backbone examples

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Amino-Liva Amino-Liva

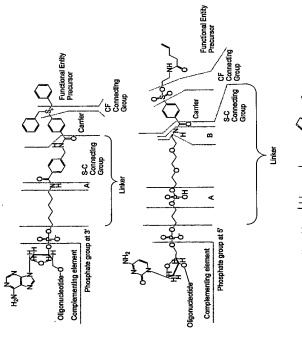
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Cena OPP-BH₃: Bormophosphates

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PCT/DK03/00176

Figure 5.



CF Connecting Attached to 5-position of pyrimidine type base S-C Connecting Group Inkar Comple menting element

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> English English

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(54) Title: A BUILDING BLOCK FORMING A C.C. OR A C.HETIERO ATOM BOND UPONREACTION
(54) Title: A BUILDING BLOCK FORMING A C.C. OR A C.HETIERO ATOM BOND UPONREACTION
(57) Abstract: A building block having the dual capabilities of transferring genetic information and functional entity precursor to a C recipient reactive group is disposed. The building block may be used in the generation of a single complex of different complexes, wherein the complex comprises are necoded molecule linked to an encoding element. Libraries of complexes are useful in the quest for pharmaceutically active compounds.

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A. CLASS	A. CLASSIFICATION OF SUBJECT MATTER IPC 7 CO7H21/00	
According	According to International Patent Classification (IPC) or to both national classification and IPC	
B. FIELDS	B. FIELDS SEARCHED	
IPC 7	Minimum obcounsations searched (classification system hollowed by classification symbots) ${ m IPC}~7~{ m CO7H}$	
Documenta	Documentation eaarched other than minimum documentation to the extent that such documents are included in the fields searched	ded in the flakts searched
Electronic o	Electronic data base consulted during the international search (name of date base and, where practical, search terms used) FPD—Internal : WPI Data	search terms used)
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C. DOCUM	C. DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with Indication, where appropriate, of the relevant passages	Relevant to claim No.
⋖	WO 00 02895 A (THOMPSON ANDREW HUGIN ;BRAX GROUP LTD (GB); SCHMIDT GUENTER (GB);) 20 January 2000 (2000-01-20) the whole document	г
۷	WALDER J A ET AL: "COMPLEMENTARY CARRIER PEPTIDE SYNTHESIS: GENERAL STRATEGY AND IMPLICATIONS FOR PREBIOTIC ORIGIN OF PRETIDE SYNTHESIS." PROCEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA, NATIONAL ACADEMY OF SCIENCE WASHINGTON, US, vol. 76, no. 1, January 1979 (1979-01), pages 51-55, KRO00857351 ISSN: 0027-8424 the whole document	50
	-/-	
×	Further documents are listed in the continuation of box C. X Patent lamby	Patent family members are listed in annex.
* Special ca	Special categories of clied documents: You can provide the property client or professy client or profesy client	ster document published after the international filing date or princity date and in conflict with the application but date in understand the principle or theory underlying the
"E" earler docr filing date "L" document v which is c	*	extraution districts relevance; the claimed invention document of particular relevance; the claimed invention of conditional to livrolve on threships stop when the document is taken alone document of particular relevance; the claimed invention
O docum other		cannot be considered to involve an inventive step when the document is combined with one or more other such docu- ments, such combination being obvious to a person skillad in the art.

P document published prior to the international filing date but later than the priority date claimed

Date of mailing of the International search report '&' document mamber of the same patent family de Nooy, A 06/10/2003 Authorized officer Name and mailing address of the ISA

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Far. (+31–70) 340–3016 Date of the actual completion of the International search 19 September 2003

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page 1 of 2

INTERNATIONAL SEARCH REPORT

Internatio pplication No PCT/DK 03/00176

20 BRUICK R K ET AL: "TEMPLATE-DIRECTED LIGATION OF PEPTIDES TO OLIGONUCLEOTIDES" CHEMISTRY AND BIOLOGY, CURRENT BIOLOGY, LONDON, 68, vol. 3, no. 1, January 1996 (1996-01), pages 49-56, XP000856876 ISSN: 1074-5521 the whole document C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT
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INTERNATIONAL SEARCH REPORT

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rhis international Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	owing reasons:
. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	
. X claims Nos.:	ils to such
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	f Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	
This International Searching Authority found multiple inventions in this international application, as follows:	
1. As all required additional search fees were timely peid by the applicant, this international Search Report covers all esearchable claims.	rrs ell
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invitis payment of any additional fee.	dta payment
 As only some of the required additional search fees were timely paid by the applicant, this international Search Report covers only those claims for which fees were paid, specifically claims Nos.: 	ch Report
4. Search Report is required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims, it is covered by claims Nos.:	Reportis
Remark on Protest The additional search fees were accompanied by the applicant's protest No protest accompanied the payment of additional search fees.	plicant's protest. ch lees.

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International Application No. PCT/DK 03 &0176